Comparison of PCR Methods for Onchocerca volvulus Detection in Skin Snip Biopsies from the Tshopo Province, Democratic Republic of the Congo

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Abstract. Defining the optimal diagnostic tools for evaluating onchocerciasis elimination efforts in areas co-endemic

INTRODUCTION

Human onchocerciasis, or river blindness, is a neglected tropical disease caused by infection with the filarial parasite, Onchocerca volvulus, and is transmitted by Simulium blackflies that breed near fast-flowing streams or rivers. An estimated 186 million people live at risk of infection.1 Onchocerciasis can cause severe itching, disfiguring skin lesions, and eye disease, which can lead to blindness.2 Mass drug administration (MDA) of the microfilaricide, ivermectin (IVM),3 administered annually for 10–15 or more years is necessary to interrupt transmission in endemic areas.4,5 Several countries in Africa have targeted elimination of the disease by April 2025, although recent modeling suggests that 2040 might be a more realistic goal for regional elimination.6

Successful elimination of onchocerciasis requires proper evaluation protocols to determine when MDA can be stopped. Laboratory tools that have been used for onchocerciasis include evaluation of skin snip biopsies (skin snips) either by microscopy or PCR testing.7–12 Detection of antibody specific for OV-16, an O. volvulus–specific antigen,13,14 and blackfly surveillance using a pool screen PCR assay targeting a repeated sequence of roughly 150 base pairs (O-150 repeat) specific for O. volvulus followed by ELISA detection (O150-PCR ELISA).15 The fact that millions of people at risk for onchocerciasis live in regions in Africa co-endemic for Loa loa or Mansonia spp., poses challenges to diagnostic testing for onchocerciasis.16–19 Mansonia streptocerca microfilariae (MF) typically migrate through the skin,20 whereas MF from both L. loa and M. perstans typically circulate within blood and not skin.18,21 The presence of MF or DNA from other filarial species in skin snips could potentially lead to false positives by microscopy or PCR methods that are not species-specific for O. volvulus.12 With the exception of a study by Wilson et al.,22 in Senegal, diagnostic performance of OV-16–based immuno-assays has been assessed mostly in populations that are not co-endemic with other filaria species.13,14,23–27 Because cross-reactivity may occur, diagnostic tools intended for onchocerciasis stop–MDA assessments need to be evaluated within several epidemiological contexts including those co-endemic for other filariae.

The 2016 WHO guidelines for stopping MDA require measuring O. volvulus prevalence in the blackfly vector using pool screen O150-PCR ELISA and prevalence in at-risk children younger than 10 years using OV-16 serology.28 Skin snip microscopy has insufficient sensitivity to be used for stopping decisions,11,12,29 but skin snip PCR can be used in some circumstances. If fewer than 10 children are found to have positive OV-16 serology, those children may be evaluated using skin snip PCR. If all children tested have negative PCR results and continue to be negative after not receiving IVM for 1 year, the evaluation areas may stop MDA, assuming the blackfly criterion has been met.28 In this situation, skin snip PCR is used to exclude patent infection in a low-prevalence setting and, thus, test sensitivity needs to be maximized. However, these guidelines do not identify which PCR method
should be implemented for this type of testing.26 This is a new procedure and to date, fewer than 10 OV-16–positive children have been tested by skin snip PCR using the O150-PCR ELISA used for blackfly vector surveillance (T. R. Unnasch, personal communication, November 2017). Because of a lack of studies that have directly compared the performance of the O150-PCR ELISA with other available PCR-based methods, there is no published evidence that the O150-PCR ELISA is the optimal PCR method for testing skin snips collected from OV-16–positive children.

Several PCR-based methods have been developed for detecting *O. volvulus*, including loop-mediated isothermal amplification,30 conventional PCR targeting the repeat O-150 region,11–10 and the O150-PCR ELISA used for blackfly vector surveillance.15 Within the last few years, real-time PCR (qPCR) methods have been developed for *O. volvulus* detection. Lloyd et al.11 described a TaqMan® (Life Technologies, Grand Island, NY) qPCR assay that targets the O-150 repeat region (qPCR-O150) and a modified conventional O150-PCR method. In addition, our group previously described the optimization of a single-reaction pan-filarial qPCR with melt curve analysis (qPCR-MCA) tool that targets a region within the first internal transcribed spacer (ITS1) of the ribosomal DNA (rDNA) and the flanking 5.8S tract12 using skin snips from sites in Ethiopia (Jimma Zone) and Uganda (Kitgum and Lamwo districts) that were not endemic for either *L. loa* or *M. perstans* (P. Cantey and V. Cama, manuscript in preparation).

Molecular tools for *O. volvulus* detection, including both the qPCR-O150 and qPCR-MCA assays, have shown increased sensitivity compared with parasitological detection by microscopy. However, varying sensitivities have been reported for different assays, perhaps, because of being tested in different epidemiological contexts. Therefore, it will be informative to determine the performance characteristics of qPCR methods and to conduct a side-by-side comparison of the O150-PCR ELISA and the two recently developed qPCR assays (qPCR-O150 or qPCR-MCA). This work will provide information on their relative sensitivity and specificity for detecting *O. volvulus* DNA in skin snips.

Thiele et al.12 demonstrated in a subset of samples (*N* = 248) that the qPCR-MCA had greater sensitivity than the O150-PCR ELISA, but the qPCR-O150 assay has not been compared with O150-PCR ELISA.11 The qPCR-MCA tool broadly amplifies the ITS1-5.8S region from multiple filarial species, relying on MCA for species categorization.12 Because of the way this tool was designed and because it was originally tested in two regions where *L. loa* and *M. perstans* were not endemic, an additional objective of this study was to further validate this tool with samples from settings that were co-endemic for *L. loa* and *M. perstans*.

The qPCR-O150 amplifies a repeat sequence,11 which has been reported to improve PCR detection; therefore, it is plausible that the qPCR-O150 might have increased sensitivity compared with the qPCR-MCA. A direct diagnostic comparison between qPCR-MCA, qPCR-O150, and O150-PCR ELISA was needed to determine which method had the greatest sensitivity and specificity and would be best suited for testing skin biopsies of OV-16–positive children.

The goals of this study were to 1) directly compare the performance of three PCR methods for *O. volvulus* detection—qPCR-O150, qPCR-MCA, and O150-PCR ELISA and 2) evaluate the qPCR-MCA tool in a setting where mixed filarial infections occurred.

**MATERIALS AND METHODS**

**Study site and sample collection.** Specimens were collected as part of a broader study designed to support the evaluation of several different diagnostic tools for onchocerciasis and other filarial infections. Samples for the evaluations described here came from the study site in the community of Banalia, Tshopo Province, Democratic Republic of the Congo (DRC). This site is co-endemic for onchocerciasis, loiasis, and Mansonellosis. Only three prior MDA distributions had occurred in this study area with the last one about 11 months before sample collection. A convenience sample of 500 participants ≥ 5 years of age who had resided within the study site for ≥ 10 years or since birth was enrolled. Consent to participate was obtained from participants aged 18 or older and from the parents of children under the age of 18 years; assent was obtained from children between the ages of 7 and 18 years. Two skin snips per participant were collected from the iliac crest and daytime blood samples were collected by venipuncture. The protocol was approved by the institutional review boards of the U.S. Centers for Disease Control and Prevention (CDC [protocol number 6196]) and the Kinshasa School of Public Health, DRC (approval number ESP/CE/032/14).

**Parasitological detection by microscopy.** Microscopic evaluation of the skin snips for MF from 471 study participants was performed as previously described using a light microscope at ×200–400 magnification.12 All samples diagnosed to have MF were verified by a second microscopist. After emergence of MF from the skin snips and visualization by microscopy, the skin snips were then preserved in 400 μL of RNAlater® (Life Technologies) and stored at –80°C until PCR testing was performed at the CDC laboratories in Atlanta, GA. The presence or absence of *L. loa* and *M. perstans* infection was assessed by microscopic evaluation of Giemsa-stained daytime thick blood smears on-site in DRC.

**DNA extraction from skin snips.** The two skin snips from each participant, which had been preserved after MF emergence and microscopic evaluation, were pooled into one DNA extraction. Genomic DNA was extracted using the QIAamp DNA Investigator Kit (Qiagen, Valencia, CA) as previously described.12 DNA quantity and quality were assessed using a NanoDrop ND-2000 spectrophotometer (Thermo Scientific, Waltham, MA).

**PCR assays.** PCR assays used in this study were performed as previously described.11,12,15,31,32 Primer and probe sequences are detailed in Supplemental Table 1. The limit of detection (LOD) for the three PCR assays for *O. volvulus* was determined and compared using serial 10-fold dilutions of adult *O. volvulus* gDNA that had been quantified using a NanoDrop ND-2000 spectrophotometer (Thermo Scientific). qPCR-O150. The qPCR assay targets the O-150 repeat region of *O. volvulus* and was performed as previously described.11 Non-endemic *Homo sapiens* gDNA and a nontemplate reaction were used as negative controls; all assays were run in duplicate. Samples were considered positive if the cycle threshold (Ct) values were below 40 amplification cycles for both duplicates. Samples were retested if duplicate qPCR-O150 results were discordant.
**qPCR-MCA.** Whole gDNA representative of single infections for *O. volvulus*, *L. loa*, *Mansonella ozzardi*, *Dirofilaria immitis*, and *Brugia pahangi* was used as references for MCA. Non-endemic *H. sapiens* gDNA and a no-template reaction were used as negative controls. All assays were run in duplicate. The melting temperature (*T_m*) ranges for MCA and filariae identification were as follows: *O. volvulus* (78.8–79.35°C, SD ± 0.21), *L. loa* (77.85–78.35°C, SD ± 0.24), and *Mansonella* spp. (76.85–77.35°C, SD ± 0.24). Previous assay optimization demonstrated that *M. perstans* and *M. ozzardi* had an indistinguishable *T_m*; therefore, *M. ozzardi* gDNA was used for the identification of *Mansonella* spp. PCR amplifications were considered positive if the Ct value was < 36 and the observed *T_m* fell within the specified range for *O. volvulus*, *Mansonella* spp., or *L. loa*. Samples with the appropriate *T_m*, but with Ct values of 36–38 were considered indeterminate and reassayed, whereas samples with a Ct > 38 were considered negative. Samples were repeated if discordant results were observed between the experimental duplicates. Species-specific qPCRs for *O. volvulus*, *M. perstans*, and *L. loa* were used to validate qPCR-MCA results.

**O150-PCR ELISA.** The O150-PCR ELISA assay was performed as previously described. Briefly, 5 μL of purified gDNA was used as template for PCR amplification. Assays included one high and one low-concentration positive control and 10 no-template negative control reactions. Conventional PCR for the O150 PCR–ELISA was performed in a T–100 thermal cycler (Bio-Rad, Hercules, CA), and ELISA detection of amplicons was measured with a SpectraMax 190 reader using SoftMax Pro v5.4.1 for data capture and analyses (Molecular Devices, Downingtown, PA). Samples were classified as positive using one of two cutoff values, always selecting the higher of the two, as previously described: 1) the mean plus three standard deviations of the 10 negative controls if this value exceeded 0.1 or 2) 0.1. Samples that were initially positive were retested beginning with a new PCR reaction. Any sample with values above the cutoff in two independent PCRs was confirmed positive.

**Species-specific TaqMan qPCRs for *M. perstans* and *L. loa*.** *Mansonella perstans* was detected using a published protocol targeting the ITS1 region of the rDNA, whereas *L. loa* was detected using a species-specific TaqMan qPCR targeting the predicted ORF LLMF72 as previously described. qPCR results were considered positive if the Ct values were below 40 amplification cycles for both duplicates. Samples were retested if duplicate results were discordant.

**Sanger sequencing.** Species specificity of TaqMan qPCRs was verified by Sanger sequencing of the ITS1 region of the rDNA, in a subset of samples that was found to be either *L. loa* (N = 16) or *M. perstans* (N = 22) positive, but not mixed. DNA sequencing chromatograms were analyzed, trimmed, and assembled using Geneious version R8 (Biomatters, Inc., Auckland, New Zealand). Filariae species were determined using the National Center for Biotechnology Information’s nucleotide database via the Basic Local Alignment Search Tool’s MegaBlast algorithm (https://blast.ncbi.nlm.nih.gov). To distinguish *O. volvulus* and *O. ochengi* sequences for the five samples that were qPCR-MCA (+) but qPCR-O150 (–), whole ITS1 sequences were compared by BLAST analysis with the *O. volvulus* and *O. ochengi* genome sequences from the WormBase ParaSite (http://parasite.wormbase.org/index.html).

**Statistical analyses.** Data analysis and statistical tests were performed using Epi Info (CDC), SPSS v. 24 (IBM SPSS Statistics, Armonk, NY), and GraphPad Prism version 7 (GraphPad Software, La Jolla, CA). Diagnostic test performance was compared using the McNemar’s test for paired nominal data, and P values were corrected for multiple comparisons using the Bonferroni correction method. A two-sided *χ²* test was performed to assess the role of non-single peak melt curve profiles in relation to the presence of mixed filarial DNA. Non-normally distributed continuous variables, as determined by D’Agostino and Pearson omnibus and Shapiro–Wilks tests for normality, were analyzed using the non-parametric Spearman’s rank correlation test.

**RESULTS**

**Filarial detection by microscopy.** Samples from 189 (40.1%) study participants were positive for *O. volvulus*, whereas none were positive for *M. streptocerca* by microscopic examination of skin snips. Microscopic evaluation of daytime thick blood smear detected 104 (22.2%) individuals with *L. loa* and 257 (54.7%) individuals with *M. perstans*, whereas 83 (17.6%) were positive for both.

**Comparison of three PCR-based methods for detection of *O. volvulus*.** Limit of detection. Overall, the qPCR-O150 had the lowest LOD at 10 fg/μL genomic DNA (Table 1). Controlling for differences in the volume of extracted DNA used in each PCR protocol, qPCR-O150 had a 2.5× lower LOD than qPCR-MCA and > 100× lower LOD when compared with O150-PCR ELISA (Table 1).

<table>
<thead>
<tr>
<th>LOD</th>
<th>DNA added (fg/μL)</th>
<th>Reaction volume (μL)</th>
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</thead>
<tbody>
<tr>
<td>qPCR-O150</td>
<td>2.0 ± 0.5</td>
<td>10 ± 0.5</td>
</tr>
<tr>
<td>qPCR-MCA</td>
<td>0.5 ± 0.1</td>
<td>10 ± 0.5</td>
</tr>
<tr>
<td>O150-PCR ELISA</td>
<td>5.0 ± 1.0</td>
<td>100 ± 5.0</td>
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**Detection of *O. volvulus* DNA in skin snip biopsies.** PCR and microscopy results for *O. volvulus* detection in skin snips are presented in Table 2. The qPCR-O150 assay detected *O. volvulus* DNA in significantly more skin snips (224) than qPCR-MCA (205; *P* = 0.003) or O150-PCR ELISA (127; *P* = 0.0003). Both qPCR methods detected more *O. volvulus*-positive skin snips than microscopy; however, the O150-PCR ELISA detected fewer positive skin snips than microscopy.

The sensitivity and specificity of each assay were compared relative to qPCR-O150, which was positive in the greatest number of *O. volvulus*-positive skin snips (Table 3). The qPCR-MCA performed similarly to the comparator, with a sensitivity and specificity of 89.3% and 98.0%, respectively. The O150-PCR ELISA had a sensitivity of 56.7% and a specificity of 100%. Sequences of the five skin snips that were negative by qPCR-O150 but positive by qPCR-MCA for *O. volvulus* all had amplicons with ≥ 99% sequence identity (615/619 bases) to *O. volvulus* sequences but only 96–98% sequence identity to *O. ochengi* genome (102/105 bases) by BLAST analysis.

**Evaluation of qPCR-MCA.** There were 266/471 skin snips (56.5%) that were positive by qPCR-MCA, indicating the presence of at least one filarial species. The *T_m* and

**Table 1** Limit of detection (LOD) for three PCR-based methods for *Onchocerca volvulus* diagnosis

<table>
<thead>
<tr>
<th>Detection method</th>
<th>DNA added (fg/μL)</th>
<th>Reaction volume (μL)</th>
<th>LOD (fg/μL)</th>
<th>LOD normalized (by vol. DNA)</th>
</tr>
</thead>
<tbody>
<tr>
<td>qPCR-O150</td>
<td>2.0 ± 0.5</td>
<td>10 ± 0.5</td>
<td>10 ± 0.5</td>
<td>20 fg</td>
</tr>
<tr>
<td>qPCR-MCA</td>
<td>0.5 ± 0.1</td>
<td>10 ± 0.5</td>
<td>100 ± 5.0</td>
<td>50 fg</td>
</tr>
<tr>
<td>O150-PCR ELISA</td>
<td>5.0 ± 1.0</td>
<td>100 ± 5.0</td>
<td>1 pg/μL</td>
<td>5 pg</td>
</tr>
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</table>
dissociation curve allowed further categorization into 205 (43.5%) O. volvulus, 19 (4.0%) L. loa, 15 (3.2%) Mansonella spp., and 27 (5.7%) as non-categorized (Table 4). qPCR-MCA results were classified as non-categorized if they displayed dissociation curves that contained more than one peak, a peak with a shoulder, or a peak that was broader than expected, as shown in Figure 1, possibly indicating the presence of more than one filarial species. Of the 24 samples that were positive by qPCR-O150 but negative by qPCR-MCA for O. volvulus in Table 3, one was positive for L. loa by MCA; two were negative by MCA, and 21 were non-categorized by MCA.

The qPCR-MCA species categorizations were compared with species-specific qPCR results (Table 4). For O. volvulus, qPCR-MCA and qPCR-O150 results were concordant in 200/224 (90.9% of the skin snips. Most of the samples non-categorized by MCA were positive for O. volvulus by qPCR-O150, with more than half of these also positive for L. loa and/or M. perstans. When comparing the qPCR-MCA categorizations to species-specific qPCR results for L. loa or M. perstans, qPCR-MCA demonstrated 100% specificity; however, the relative sensitivity for detecting L. loa DNA (47.5%) and M. perstans DNA (30%) in skin snips was low.

A total of 46 (9.8%) samples had DNA from multiple filarial species detected. qPCR-MCA non-categorized skin snips (N = 27) were more likely to contain DNA from multiple filarial species—20/46 mixed infections had non-categorized results and 7/215 single infections had non-categorized results (P < 0.0001, two-sided χ²). Mixed filarial DNA, however, was detected in 26 cases when the dissociation curve was a single peak.

The frequency of the different filarial species DNA present in the skin snips defined by species-specific qPCR results is presented in Table 5. Overall, 208 (44.2%) of the skin snips were negative, 224 (47.5%) had O. volvulus, 40 (8.5%) had L. loa, and 50 (10.6%) had M. perstans. Most of the individuals with skin snips that were L. loa qPCR positive were also positive for this parasite by blood smear microscopy (35/40 or 87.5%). Likewise, a majority of individuals with skin snips that were M. perstans qPCR positive were also positive by blood smear microscopy (48/50 or 96%).

### Comparison of O150-based PCR methods

The O150-PCR ELISA was less sensitive than either of the two qPCR methods tested. Therefore, potential factors influencing reduced O150-PCR ELISA sensitivity were investigated. A total of 161/471 (34.2%) samples were initially positive by O150-PCR ELISA. However, 34/161 (21% of the initial positives) were negative on repeat testing, so the final result was negative (per protocol). Twenty-four of those 34 negatives on repeat testing (70.6%) were qPCR-O150(+). Receiver operator characteristics analysis for O150-PCR ELISA showed the optimal threshold at OD = 0.055. Compared with the standard threshold at 0.1, the use of this optimal threshold would increase the sensitivity from 56.7% to 83%, but specificity would be reduced from 100% to 90.7% (Supplemental Figure 1). Comparison of the sensitivity of the qPCR-O150 and O150-PCR ELISA methods showed decreased O150-PCR ELISA sensitivity in skin snips with lower MF loads (Figure 2A). Because the qPCR-O150 Cₘ values are semi-quantitative and highly correlated with MF load (Supplemental Figure 2; Spearman’s rank ρ = −0.74, P < 0.0001), the O150-PCR ELISA sensitivity was compared with quartiles of Cₘ values (Figure 2B); this showed a decline in O150-PCR ELISA sensitivity with higher Cₘ values.

### DISCUSSION

We directly compared the performance of three PCR-based methods for O. volvulus detection—two real-time methods (qPCR-O150 and qPCR-MCA) and one conventional PCR-based method (O150-PCR ELISA)—using skin snips from an onchocerciasis endemic region co-endemic for L. loa and M. perstans. We also evaluated the qPCR-MCA tool in a setting co-endemic for several filariae. The results show that qPCR-O150 detected O. volvulus DNA in significantly more skin snips than either of the other PCR methods tested. Notably, the O150-PCR ELISA currently in use by onchocerciasis programs detected only 56.7% of the skin snips in which O. volvulus DNA was detected by qPCR-O150. The reduced sensitivity of both tests compared with qPCR-O150 was because of differences in limits of detection among the assays. As a result, the O150-PCR ELISA was less sensitive in samples with lower counts of MF. However, specific issues with the qPCR-MCA leading to reduced sensitivity also included the misclassifications of filarial species in the cases of mixed infections.

There were 10 individuals who were microscopy positive but negative by qPCR-O150 and O150-PCR ELISA. The MF loads in these 10 samples were low (median 1.5 MF/2 snips; range 1.0–22), and we cannot exclude the possibility that PCR amplifications were negative because of our use of

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**Table 2**

<table>
<thead>
<tr>
<th>Detection method</th>
<th>Positive</th>
<th>% Positive (%)</th>
<th>P value*</th>
</tr>
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<tbody>
<tr>
<td>qPCR-O150</td>
<td>224</td>
<td>47.5</td>
<td>−</td>
</tr>
<tr>
<td>qPCR-MCA</td>
<td>205</td>
<td>43.5</td>
<td>0.003</td>
</tr>
<tr>
<td>O150-PCR ELISA</td>
<td>127</td>
<td>27.0</td>
<td>0.0003</td>
</tr>
<tr>
<td>Microscopy</td>
<td>189</td>
<td>40.1</td>
<td>0.0003</td>
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</table>

* McNemar’s test with Bonferroni correction for multiple comparisons.

**Table 3**

<table>
<thead>
<tr>
<th></th>
<th>qPCR-O150</th>
</tr>
</thead>
<tbody>
<tr>
<td>Positive</td>
<td></td>
</tr>
<tr>
<td>Negative</td>
<td></td>
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<tr>
<td>Total</td>
<td></td>
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<tr>
<td>Sensitivity</td>
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<tr>
<td>Specificity</td>
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The qPCR-MCA tool detected skin snips that displayed a $T_m$ consistent with either *O. volvulus*, *L. loa*, or *Mansonella* spp., accurately detecting 89.3% of *O. volvulus*–positive skin snips. However, this tool was unable to categorize 27 samples because of their irregular melt curve profiles. Compared with species-specific qPCRs, the qPCR-MCA had lower sensitivity for detecting *L. loa* and *M. perstans* in skin snips than for detecting *O. volvulus*. It is plausible that in an onchocerciasis hyperendemic setting such as the one tested here, the relative abundance of DNA from *M. perstans* and *L. loa*, which are not typically found in the skin, would be lower than that from *O. volvulus* DNA and, thus, might not always be detected by the MCA tool. Another point to consider is if *O. volvulus* MF loads are suppressed by multiple rounds of IVM, the ability of the qPCR-MCA to detect *O. volvulus* might decrease even further in the context of coinfections with other filariae.

The use of species-specific qPCRs was imperative to evaluate the performance of qPCR-MCA because initial assay optimization demonstrated that $T_m$ profiles could not discriminate between closely related species, limiting its discrimination to the genus level.12 qPCR-MCA alone cannot discriminate *M. streptocerca*, which typically circulates within the skin,20 from *M. perstans*. Furthermore, another potential limitation of qPCR-MCA might be its inability to distinguish between *O. volvulus* and other closely related *Onchocerca* species such as *O. ochengi* and *Onchocerca* sp. “Siisa,” which can be found in African blackflies,35–37 but we did not directly test this hypothesis. Interestingly, five skin snips that were categorized as qPCR-MCA(+) for *O. volvulus* were qPCR-O150(–), with all five demonstrating ITS1 sequence similarity to *O. volvulus*. These five qPCR-O150 false negatives might be because of suboptimal primer/probe binding. Although the qPCR-MCA identified five positive results that were not detected by the qPCR-O150, this assay nevertheless was less sensitive than qPCR-O150. A more extensive validation of qPCR-MCA for detection of filarial species within other sample types derived from blood is necessary to assess the utility of this tool within other contexts.

Despite the limitations of qPCR-MCA, its use in DRC allowed for the unexpected identification of *L. loa* and *M. perstans* DNA in skin snips samples. The presence of DNA from these two species might be more common in skin snips than previously recognized. Whether MF are present and thus could be sources of false positives by microscopy remains unclear. However, MF from both *L. loa* and *M. perstans* are not usually found circulating in the skin.18,21 Therefore, one plausible explanation for the presence of DNA of these species in skin snips could be the unintentional capture of blood during the biopsy; however, this seems unlikely given the fact that there were blood smear negative individuals that were skin snip PCR positive. Alternatively, the PCR assays might be detecting cells or circulating DNA from either *M. perstans* or *L. loa* because adults migrate through subcutaneous tissue.21

![Figure 1](image-url) Representative melt curve profiles from two different skin snip DNA samples that displayed non-single peak profiles indicative of mixed template DNA. Species-specific qPCR results for these samples indicated the presence of both *Mansonella perstans* and *Onchocerca volvulus*. The $T_m$ range for *O. volvulus* and *Mansonella* spp. (+) positive controls is indicated with checkered gray and solid gray bars, respectively. The melt curve profile from each skin snip sample is shown in black (duplicate replicates in solid vs. dashed line) (A) melt curve profile with a main peak and a secondary smaller peak. (B) Melt curve profile with two distinct peaks.
Currently, the O150-PCR ELISA test is the standard tool used for detection of *O. volvulus* in *Simulium* blackflies, and thus far, it has been used for confirmation of patent infection in less than 10 children with positive OV-16 antibody results (T. R. Unnasch, personal communication, Nov. 2017). As mentioned previously, the sensitivity of this method as tested was low and decreased as *O. volvulus* MF and DNA decreased, corroborating the observations of Thiele et al.\(^\text{12}\) As PCR will be used to exclude patent infection in children positive for OV-16 serologic test, sensitivity is very important and efforts should be made to maximize the sensitivity of the PCR used by programs. We showed that the qPCR-O150 assay would be the most appropriate test for this based on its LOD and high sensitivity.

Although qPCR technology is currently available in many onchocerciasis endemic countries, some onchocerciasis programs might not have access to qPCR but would have access to the equipment used for O150-PCR ELISA testing of blackflies. A detailed comparison of the accuracy and sustainability of qPCR versus O150-PCR ELISA would be necessary to decide which PCR method would be the best to properly inform the onchocerciasis elimination programs. In the event that qPCR-O150 could not be implemented by a program, consideration will need to be given for maximizing the sensitivity of the O150-PCR ELISA to improve its detection performance, especially when MF loads are low. Adjusting the O150-PCR ELISA OD cutoff could improve sensitivity, but at a cost to specificity. Assay sensitivity is also important for blackfly surveillance, but for blackfly testing, lower sensitivity can be circumvented to an extent by increasing the numbers of blackflies tested. This cannot be performed when testing OV-16–positive children. However, increased sensitivity with loss to specificity could be important in posttreatment surveillance of blackflies to maximize early recognition of recrudescence. Testing an additional locus such as cytochrome oxidase I may also help to overcome the potential loss to specificity.\(^\text{38}\) An oligonucleotide-based magnetic bead capture method for purification of *O. volvulus* DNA from blackfly vectors method was previously shown to increase detection sensitivity.\(^\text{15}\) Extracting DNA from skin snips using this method might further increase the sensitivity of the O150-PCR ELISA in skin snips; however, additional studies are needed to evaluate this.

### TABLE 5

<table>
<thead>
<tr>
<th>Filarial species</th>
<th>Total #</th>
<th>Total (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total positive</td>
<td>263</td>
<td>55.8</td>
</tr>
<tr>
<td><em>Onchocerca volvulus</em></td>
<td>224</td>
<td>47.5</td>
</tr>
<tr>
<td><em>Loa loa</em></td>
<td>40</td>
<td>8.5</td>
</tr>
<tr>
<td><em>Mansonella perstans</em></td>
<td>50</td>
<td>10.6</td>
</tr>
<tr>
<td>Total single infections</td>
<td>217</td>
<td>46.1</td>
</tr>
<tr>
<td><em>O. volvulus</em></td>
<td>185</td>
<td>39.3</td>
</tr>
<tr>
<td><em>L. loa</em></td>
<td>16</td>
<td>3.4</td>
</tr>
<tr>
<td><em>M. perstans</em></td>
<td>16</td>
<td>3.4</td>
</tr>
<tr>
<td>Total mixed infections</td>
<td>46</td>
<td>9.8</td>
</tr>
<tr>
<td><em>O. volvulus</em> + <em>L. loa</em></td>
<td>12</td>
<td>2.5</td>
</tr>
<tr>
<td><em>O. volvulus</em> + <em>M. perstans</em></td>
<td>22</td>
<td>4.7</td>
</tr>
<tr>
<td><em>O. volvulus</em> + <em>L. loa</em> + <em>M. perstans</em></td>
<td>5</td>
<td>1.1</td>
</tr>
<tr>
<td><em>L. loa</em> + <em>M. perstans</em></td>
<td>7</td>
<td>1.5</td>
</tr>
<tr>
<td>Negative</td>
<td>208</td>
<td>44.2</td>
</tr>
<tr>
<td>Total</td>
<td>471</td>
<td>100.0</td>
</tr>
</tbody>
</table>

*As defined by qPCR-O150.*

Even if the O150-PCR ELISA protocol for PCR-testing of skin snips could be adapted to augment sensitivity, the qPCR-O150 assay still affords several advantages over the O150-PCR ELISA method. qPCR techniques do not require any post-amplification processing, thus reducing the possibility for cross-contamination and need for extensive quality control measures. The assay time for the qPCR-O150 assay is less than 2 hours, much shorter than for O150-PCR ELISA, which can take up to 2 days. Moreover, the qPCR-O150 assay requires less equipment compared with the O150-PCR ELISA, which uses a regular PCR thermocycler, ELISA plate reader, incubators, and possibly an ELISA plate washer. The qPCR is also simpler to perform because there are fewer reagents and
protocol steps involved. This may reduce interoperator assay variability and would make it more amenable to standardization across diverse laboratory environments. These same advantages would also apply to qPCR testing of blackflies; thus, comparing the sensitivity and specificity of qPCR-O150 versus O150-PCR ELISA in blackflies could be important, particularly for posttreatment surveillance, when early detection of recrudescence will be important. It would also be worthwhile to conduct a comprehensive comparison between the two assays, both for testing of blackflies and skin snips, to determine if it would be beneficial to programatically implement the qPCR-O150 assay both for the evaluation of blackflies and skin snips.

In summary, we found the qPCR platform to be more sensitive than O150-PCR ELISA. Increasing the sensitivity of the currently available PCR technique or using real-time PCR technology, if logistically and technically feasible, should be considered. As increasing the sensitivity for O. volvulus detection in blackflies could enhance earlier detection of recrudescence, an evaluation of these PCR methods in blackflies could be an important next step.

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